AMENDMENTS TO THE CLAIMS

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- 1. (currently amended) Method for selective separation by density of each constituent of a mix of organic synthetic materials and particularly comminuted waste materials to be reused by recycling, stable within a precision level of ± 0.0005 about a density level [["]]ds[["]] chosen as the density separation threshold equal to at least 1, and consisting of comprising separating them by density difference, in an aqueous suspension of an appropriate quantity of powder particles dispersed in a sufficient quantity to create the density level [["]]ds[["]] chosen as the separation threshold of at least one of the constituents of the mix of fragmentated synthetic organic materials to be selectively separated, characterised in that wherein the separative suspension is formed from:
 - a) solid powder particles with a size grading cutoff not more than 30 μm, these solid powder particles thus sized being dispersed in the aqueous phase in a sufficient quantity to reach the chosen density threshold [["]]ds[["]],
 - b) a water soluble stabiliser to stabilise the rheological and invariance characteristics of the density [["]]ds[["]] with a precision of the said density [["]]ds[["]] of the suspension of solid powder particles equal to ±0.0005.
- 2. (currently amended) Method according to claim 1, characterised in that wherein the solid powder particles preferably have a size grading cut-off not more than 20 μ m and even better not more than 5 μ m, these solid powder particles thus sized being dispersed in the aqueous phase in a sufficient quantity to create the chosen density threshold [["]]ds[["]].
- 3. (currently amended) Method according to claim 1, wherein or 2, characterised in that the powder particles are of natural origin and are at least one of chosen from the group of powder mineral materials consisting of clays, belonging to families consisting of the group composed of kaolinites, comprising kaolinite, dickite, halloysite, disordered kaolinites, serpentines, the micas group and particularly muscovite, biotite, paragonite, pyrophyllite and talc, illites and glauconite, the montmorillonites group and particularly beidellite, stevensite, saponite, hectorite; the chlorites group; the vermiculites group; the interstratified clays group for which the unit structure

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is a combination of the previous groups; the fibrous clays group, and particularly attapulgite

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(palygorskite), sepiolite; the group consisting of calcium carbonate (calcite), magnesium

carbonate, dolomite (double-calcium and magnesium carbonate), dehydrated calcium sulphate

(gypsum), barium sulphate, talc, alumina, silica, titanium dioxide, zirconia, taken alone or mixed.

4. (currently amended) Method according to claim 1, wherein or 2, characterised in that the

powder particles are of synthetic origin and are chosen from among the group consisting of glass

powders, precipitated calcium carbonate, metallic powders, taken alone or mixed.

5. (currently amended) Method according to at least one of claim[[s]] 1 to 4, characterised

in that wherein the median diametric dimension of the powder particles is limited to 5 µm and

preferably between 1- µm and 0.005 µm.

6. (currently amended) Method according to one of claim[[s]] 1 to-5, characterised in that

wherein the water soluble agent for stabilisation of rheological characteristics and to maintain the

apparent density [["]]ds[["]] of the suspension of solid powder particles, is chosen from the

group composed of phosphates and polyphosphates, alkylphosphate esters, alkylphosphonate,

alkylsulfate, alkylsulfonate esters, lignin, lignosulfonates in the form of calcium, sodium, iron,

chromium, iron and chromium salts, copolymers of maleic anhydride and sulfonic styrene acid,

substituted, neutralised, esterified or non-esterified copolymers of methylacrylamide and

(methyl) acrylic acid, copolymers of sulfonic methylacrylamido-alkyl acid and (methyl)

acrylamide acid, water soluble polymers of acrylic acid used in acid form of possibly wholly or

partly neutralised by alkaline and/or alkali earth agents, by amines salified by single-valent

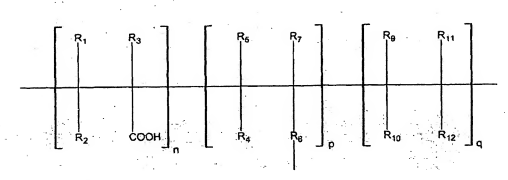
and/or polyvalent ions, and/or esterified, or by water soluble acrylic copolymers with

phosphated, phosphonated, sulphated or sulfonated functions.

7. (currently amended) Method according to claim 6, characterised in that wherein the

water soluble agent is preferably chosen from among water soluble acrylic copolymers satisfying

the general formula:



in which:

 $\sqrt{2}$ is a phosphate, phosphonate, sulphate, sulfonate type pattern with at least one free acid function,

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- $\sqrt{}$ n is equal to a value of between 0 and 95,
- $\sqrt{}$ p is equal to a value of between 95 and 5,
- $\sqrt{\ }$ q is equal to a value of between 0 and 95,
- $\sqrt{ }$ the total value of n + p + q is equal to 100,
- $\sqrt{R_1}$ and R_2 are hydrogen simultaneously, or one is hydrogen while the other is a carboxylic function that may or may not be esterified by an alcohol in C_1 to C_{12} ,
- $\sqrt{R_3}$ is hydrogen or an alkyl radical in C_1 to C_{12} ,
- $\sqrt{R_4}$ and R_5 are simultaneously or separately hydrogen or an alkyl radical in C_1 to C_{12} , a substituted or non-substituted aryl, a carboxylic function that may or may not be esterified by an alcohol in C_1 to C_{12} ,
- $\sqrt{R_6}$ is a radical that sets up the link between the pattern \mathbb{Z} and the polymeric chain, this radical R_6 possibly being an alkylene with formula CH_{2r} in which r may be equal to values in the interval 1 to 12, an oxide or polyoxide of alkylene with formula R_8O_s in which R_8 is an alkylene in C_1 to C_4 and s may be any value from 1 to 30, or a combination of the two formulas CH_{2r} and R_8O_s ,
- $\sqrt{R_7}$ can be hydrogen or an alkyl radical in C_1 to C_{12} ,

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- $\sqrt{R_9}$ and R_{10} are simultaneously hydrogen, or one is hydrogen and the other is a carboxylic group, an ester in C_1 to C_{12} , an alkyl in C_1 to C_{12} , an aryl in C_5 or C_6 or an alkylaryl,
- $\sqrt{R_{11}}$ may be hydrogen, a carboxylic group, an alkyl in C_1 to C_3 or a halogen,
- $\sqrt{R_{12}}$ may be an ester in C_1 to C_{12} , a substituted or non-substituted amide, an alkyl in C_1 to C_{12} , an aryl in C_5 or C_6 , an alkylaryl, a halogen, a carboxylic group or an alkyl or aryl phosphated, phosphonated, sulphated, sulfonated group.
- 8. (currently amended) Method according to claim 7, characterised in that wherein, in pattern Z, the non-free acid functions are occupied by a cation, an ammonium group, an amine, an alkyl in C_1 to C_3 , an aryl in C_3 to C_6 substituted or not, an alkylaryl, an ester in C_1 to C_{12} and preferably in C_1 to C_2 , or a substituted amide.
- 9. (currently amended) Method according to claim 8, characterised in that wherein the total value of n + p + q, n = 0, when q > 0 and q = 0 when n > 0.
- 10. (currently amended) Method according to claim 8, characterised in that wherein, in patterns R_1 and R_2 , the alcohol esterifying the carboxylic function is preferably C_1 to C_4 .
- 11. (currently amended) Method according to claim 9, characterised in that wherein, in pattern R_3 , the alkyl radical is preferably in C_1 to C_4 .
- 12. (currently amended) Method according to claim 8, characterised in that wherein, in patterns R_4 and R_5 , the alkyl radical is preferably in C_1 to C_4 .
- 13. (currently amended) Method according to claim 8, wherein characterised in that, in patterns R_4 and R_5 , the alcohol esterifying the carboxylic function is preferably in C_1 to C_4 .

14. (currently amended) Method according to claim 8, characterised in that wherein, in pattern R_7 , the alkyl radical is preferably in C_1 to C_4 .

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- 15. (currently amended) Method according to claim 8, characterised in that wherein, in patterns R_5 and R_{10} , the ester is preferably in C_1 to C_3 .
- 16. (currently amended) Method according to claim 7, characterised in that wherein, in patterns R_9 and R_{10} , the alkyl is preferably in C_1 to C_3 .
- 17. (currently amended) Method according to claim 7, characterised in that wherein, in pattern R_{12} , the ester is preferably in C_1 to C_5 .
- 18. (currently amended) Method according to claim 7, characterised in that wherein, in pattern R_{12} , the alkyl is preferably in C_1 to C_3 .
- 19. (currently amended) Method according to claim 7, eharacterised in that wherein the molecular mass \underline{M}_{w} of water soluble acrylic copolymers forming the stabilisation agent is between 5000 and 100 000.
- 20. (currently amended) Method according to claim 7, characterised in that wherein the water soluble acrylic copolymers forming the stabilisation agent are at least partially neutralised, using a neutralisation agent chosen from the group composed of sodium, potassium, ammonium, calcium, magnesium hydroxides, and primary, secondary or tertiary, aliphatic and / or cyclic amines such as mono, di, tri ethanolamines, mono and diethylamines, cyclohexylamine, methylcyclohexylamine, etc.
- 21. (currently amended) Method according to at least one of claim[[s]] 1 to 20, characterised in that wherein the quantity by weight of the water soluble stabilisation agent is expressed as the dry weight of the said agent as a percentage of the dry weight of powder particles put in

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suspension, and is between 0.02% and 5% [[,]] and preferably between 0.1% and 2% dry weight

based on [[of]] the dry weight of the powder particles.

22. (currently amended) Method according to at least one of claim[[s]] 1 to 21, characterised

in that wherein the aqueous phase has a conductivity of not more than 50 ms, and preferably

between 0.2 ms and 40 ms.

23. (currently amended) Method according to at least one of claim[[s]] 1 to 22, characterised

in that it wherein said method is carried out in at least one hydraulic separator.

24. (currently amended) Method according to claim 23, characterised in that wherein, when

the said method is carried out in a single hydraulic separator, the chosen density [["]]ds[["]]

varies:

- in increasing order, by controlled addition of the defined powder particles and the

water soluble stabilisation agent in the suspension present in the said hydraulic

separator, until the new chosen threshold density [["]]ds[["]] is obtained,

in decreasing order, by adding water until obtaining the new chosen threshold

density [["]]ds[["]].

25. (currently amended) Method according to claim 24, eharacterised in that wherein the

density of the separative stable suspension is varied, to increase it or to decrease it, while

maintaining mechanical stirring of the dense aqueous medium being corrected and / or while

recirculating the dense medium by drawing off from the bottom of the hydraulic separator and

reinjection at the top of the said separator.

26. (currently amended) Method according to claim 23, characterised in that wherein if the

said method is performed in several hydraulic separators, the various separators are placed one

after the other, using a cascade system operating with stable suspensions each having a precise

density threshold [["]]ds[["]], in an increasing or decreasing order of the densities.

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27. (currently amended) Method according to any one of claim[[s]] 23 to 26, characterised in

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that wherein the threshold density [["]]ds[["]] of the precise aqueous separative suspension is

continuously controlled by appropriate measurement means and is corrected as soon as a drift is

detected.

28. (currently amended) Method according to claim 27, characterised in that wherein the

density of the dense medium contained in each hydraulic separator is controlled using two

solenoid valves each opening onto two circuits connected to two reservoirs, one of the reservoirs

containing a concentrated [["]]mother[["]] suspension of powder particles, stabilised with a water

soluble stabilisation agent so that a determined quantity of the mother suspension can be added

to make an upward correction to any downward drift in the density of the dense precise

separation medium, the other tank containing water so that a determined quantity of water can be

added to make a downward correction to any upward drift in the density of the dense precise

separation medium.

29. (currently amended) Method according to claim 28, characterised in that wherein the

density of the dense medium in each hydraulic separator is measured continuously by

appropriate measurement instruments that trigger opening of one of the solenoid valves and then

close it when the threshold density [["]]ds[["]] is reached.

30. (currently amended) Method according to any one of claim[[s]] 23 to 29, characterised in

that wherein each suspension may be recirculated in each hydraulic separator by drawing off the

said suspension at the bottom of the said separator and reinjecting it into the upper part.

31. (currently amended) A method of claim 1, wherein said method is Use of the method

according to at least one of claims 1 to 30; for selectively separating mixed polymer

materials[[,]] and particularly waste materials originating from the destruction of cars and /or

durable consumer products at the end of their-lives.